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GRAFT COPOLYMERS AND IMPACT-RESISTANT FLAME-RETARDANT RESIN (54)COMPOSITIONS CONTAINING THE SAME

(57) The present invention provides a polyorganosiloxane-containing graft copolymer

which is obtainable by polymerizing 0.5 to 10 parts by weight of a vinyl monomer (B) comprising 100 to 50% by weight of a polyfunctional monomer (b-1) containing two or more polymerizable unsaturated bonds in the presence of 40 to 90 parts of polyorganosiloxane particles,

followed by further polymerizing 5 to 50 parts by weight of a vinyl monomer (C):

A polyorganosiloxane-containing graft copolymer which is obtainable by polymerizing 0 to 10 parts by weight of a vinyl monomer (B) comprising 100 to 50% by weight of a polyfunctional monomer containing two or more polymerizable unsaturated bonds in the presence of 30 to 95 parts of a polyorganosiloxane in a latex form as obtained by seed polymerization using, as a seed polymer, a hydrophilic polymer capable of swelling in the corresponding organosiloxane,

followed by further polymerizing 5 to 70 parts by weight of a vinyl monomer (C);

a flame retardant which comprises said copolymer: and

a resin composition which comprises said retardant and a thermoplastic resin.

Description

TECHNICAL FIELD

5 [0001] The present invention relates to polyoganosilioxane-containing graft copolymers and impact-resistant, flame-retardant resin compositions containing the same.

BACKGROUND ART

- 10 [0002] Owing to their good impact resistance, heat resistance and electric characteristics, among others, thermoplastic resins, in particular polycarbonally resins, are widely used as materials of electric and electronic parts, OA (office
 automation) apparatus and instruments, and household utensify, or as building materials. Polycarbonate resins, though
 higher in flame retardancy as compared with polystyrene and other resins, are required to be highly flame retardants
 retardioular in such fields as electric and electronic parts. OA apparatus and instruments and the like and, therefore,
 particular in such fields as electric and electronic parts. OA apparatus and instruments and the like and, therefore,
 particular in such fields as electric and electronic parts. OA apparatus and instruments and the like and, therefore,
 attempts have been made to improve their flame retardancy by adding various flame retardants. Thus, for instance,
 the addition of organohalogen compounds or organophosphorus compounds have a problem from the toxicity viewpoint. In particular, it is a drawback of organophalogen compounds they generate a corrosive gas upon combustion thereof. Thus,
 the delivers of the property of the property
- the demand for halogen-free and phosphorus-free flame retardants has been increasing in recent years.

 The utilization of polyorganosiloxane compounds (also called silicones) as halogen-free and phosphorus-free [0003]. The utilization of polyorganosiloxane compounds (also called silicones) as halogen-free and phosphorus-free [0004]. Japanese Kokal Publication Sho-54-36856 describes that kneading of a monorganopolysiloxane-based silicone resin with a non-silicone polymer gives a flame retardant resin.

 [0004] Japanese Kohyo Publication Heli-3-48947 describes that a mixture of a silicone resin and a saft of a metal of the control of the control
- the group IIA provides thermoplastic resins with flame retardancy.

 [0005] Japanese Kokai Publication Hel-E-113712 describes a method of producing flame retardant resin compositions which comprises Kokai Publication Hel-E-113712 describes a method of producing flame retardant resin compositions which comprises dispersing a silicone resin propared by blending 100 parts by weight of a polyorganositoxane with 10 to 150 parts by weight of a silical filter in thermoplastic resins.
- [0006] Japenese Kokai Publication Hei-10-139964 describes that flame retardant resin compositions are obtained by adding a solvent-soluble silicone resin having a weight average molecular weight of not less than 10,000 but not one than 270,000 to an aromatic ring-containing has become resin average molecular weight of not less than 10,000 but not one than 270,000 to an aromatic ring-containing has because the flame of the containing the provided numbers are indeed effective in providing flame.
 - [0007] However, the silicone resins described in the above-cited publications are indeed effective in providing flame retardancy but their effects are still unsatisfactory. When the addition level is increased to fill up the shortage, a problem arises that the impact resistance of the resin composition decreases, making it difficult to obtain flame retardant resin composition.
- composition balanced between flame retardancy and impact resistance.

 [008] Japanese fokair Publication 2000-17029 describes that when a composite rubber-based flame retardant pro[008] Japanese fokair Publication 2000-17029 describes that when a composite rubber based of a polyographic state of the properties of th
- [0009] Japanese Kokai Publication 2000-226420 describes that flame retardant resin compositions can be obtained by incorporating a polyorganosiloxane-based flame retardant produced by grafting a vinyl monomer onto composite particles consisting of an aromatic group-containing polyorganosiloxane and a vinyl polymer in thermoplastic resins. [0010] Japanese Kokai Publication 2000-264935 describes that flame retardant resin compositions can be obtained by incorporating, in thermoplastic resins, a Polyorganosiloxane-containing graft copolymer prepared by graft copolymer prepared by graft copolymer prepared by graft copolymer programs.
- merization of a vinyl monomer onto polyorganosiloxane particles not larger than 0.2 µm in size

 [0011] The flame retardant resin compositions described in the above-cited Japanese Kokai Publication 2000-216420 and Japanese Kokai Publication 2000-226420 and Japanese Kokai Publication 2000-226420 and Japanese Kokai Publication 2000-226420 and Japanese Hotal Publication 2000-226430 and Japanese Hotal Publication 2000-226430 and Japanese Kokai Publication

50 SUMMARY OF THE INVENTION

[0012] It is an object of the present invention to provide a polyorganosiloxane-containing graft copolymer utilizable as a halogen-free and phosphorus-free flame retardant and excellent in flame retardancy and impact resistance improving effects as well as a flame-retardant resin composition excellent in flame retardancy and impact resistance using the order to the composition excellent in flame retardancy and impact resistance using the order to the composition excellent in flame retardancy and impact resistance using the composition excellent in flame retardancy and impact resistance.

using the graft oppolymer mentioned above.

[0013] The present inventors made intensive investigations concerning the above subject and, as a result, found that a specific polyorganosiloxane-containing graft copolymer is excellent in flame retardancy and impact resistance that a specific polyorganosiloxane-containing graft copolymer is excellent in flame retardancy and impact resistance can improving effects and that a flame-retardant resin composition excellent in flame retardancy and impact resistance can improving effects and that a flame-retardant resin composition excellent in flame retardancy and impact resistance can improving effects and that a flame-retardant resin composition excellent in flame retardancy and impact resistance.

be obtained by incorporating the polyorganosiloxane-containing graft copolymer in a thermoplastic resin. Based on such findings, the present invention has now been completed.

[0014] Thus, in accordance with a first aspect thereof, the present invention relates to:

- a polyorganosiloxane-containing graft copolymer which is obtainable by polymenzing 0.5 to 10 parts (parts by weight; hereinafter the same shall apply) of a vinyl monomer (B) comprising 100 to 50% (% by weight; hereinafter the same shall apply) of a polyfunctional monomer (b-1) containing two or more polymerizable unsaturated bonds and 0 to 50% of another copolymenzable monomer (b-2), in the presence of 40 to 90 parts of polyorganosiloxane
- followed by further polymerization of 5 to 50 parts of a vinyl monomer (C) (the sum of (A1), (B) and (C) being 100 10 parts] (Claim 1);

the polyorganosiloxane-containing graft copolymer according to Claim 1,

- wherein the polyorganosiloxane particles (A1) have a volume average particle diameter of 0.008 to 0.6 μm (Claim continuous). 15
 - 2): the polyorganosiloxane-containing graft copolymer according to Claim 1 or 2,
 - wherein the vinyl monomer (C) gives a polymer thereof having a solubility parameter of 9.15 to 10.15 (cal/cm³) 1/2 (Claim 3);
 - the polyorganosiloxane-containing graft copolymer according to any of Claims 1 to 3,
 - wherein the polyorganosiloxane particles (A1) are in a latex form (Claim 4);
 - the polyorganosiloxane-containing graft copolymer according to any of Claims 1 to 4, wherein the vinyl monomer (C) comprises at least one monomer selected from the group consisting of aromatic vinyl monomers, vinyl cyanide monomers, (meth)acrylate ester monomers and carboxyl group-containing vinyl monomers (meth)acrylate ester monomers and carboxyl group-containing vinyl monomers. omers (Claim 5):
- 25 a flame retardant

- which comprises a polyorganosiloxane-containing graft copolymer according to Claim 1 (Claim 6);
- which comprises 0.1 to 30 parts, per 100 parts of a thermoplastic resin, of a flame retardant according to Claim 6 as incorporated in the thermoplastic resin (Claim 7); and 30
 - the flame retardant resin composition according to Claim 7,
 - wherein the thermoplastic resin is a polycarbonate resin (Claim 8).
 - [0015] In accordance with a second aspect thereof, the invention relates to:
- a polyorganosiloxane-containing graft copolymer which is obtainable by polymerizing 0 to 10 parts by weight of a vinyl monomer (B) comprising 100 to 50% by weight of a polyfunctional monomer (b-1) containing two or more 35 polymerizable monomer (b-2), in the presence of the company of the ence of 30 to 95 parts by weight (as solid content) of a polyorganosiloxane (A2) in a latex form obtainable by seed polymerization using, as a seed polymer, a hydrophilic polymer capable of swelling in the corresponding organosi-
- followed by further polymerizing 5 to 70 parts by weight of a vinyl monomer (C) (the sum of (A2), (B) and (C) being 100 parts] (Claim 9);
 - the polyorganosiloxane-containing graft copolymer according to Claim 9,
- wherein the seed polymer has such a degree of hydrophilicity that the extraction rate of water-soluble components in dry seed polymer has such a degree onlymer has such a degree onlymer is 10 to 100% by weight, as determined after adding water, in an amount of 20 weight, as determined after adding water and a weight of 20 weight. of the seed polymer in a dry state, to the dry seed polymer, followed by stirring at 23 °C for 1 hour and
 - wherein the seed polymer shows such a degree of swelling in the organosiloxane that the rate of swelling by volume as determined from the ratio between the latex particle diameter after stirring and the latex particle diameter before stirring in an amount 50 times by weight that of the dry seed before stirring is 3 to 50 times after adding the organosiloxane, in an amount 50 times by weight that of the dry seed polymer, to the post of the polymer to the polymer t
 - polymer, to the seed polymer latex, followed by stirring at 23 °C for 1 hour (Claim 10); the polyorganosiloxane-containing graft copolymer according to Claim 9,
 - wherein the seed polymer has such a degree of hydrophilicity that the extraction rate of water-soluble components in dry seed polymer is 50 to 100% by weight. as determined after adding water, in an amount of 20 weight-times that of the seed polymer is 50 to 100% by weight. of the seed polymer in a dry state, to the dry seed polymer, followed by stirring at 23 °C for 1 hour, and
 - wherein the seed polymer shows such a degree of swelling in the organosiloxane that the rate of swelling by volume as determined from the ratio between the latex particle diameter after stirring and the latex particle diameter before stirring is 3 to 15 times after adding the organosiloxane, in an amount 50 weight-times that of the dry seed polymer. (blue next) 15 times after adding the organosiloxane, in an amount 50 weight-times that of the dry seed polymer. polymer, to the seed polymer latex, followed by stirring at 23 °C for 1 hour (Claim 11);

[0016] The polyorganosiloxane-containing graft copolymer according to any of Claims 9 to 11,

wherein the vinyl monomer (C) comprises at least one monomer selected from the group consisting of aromatic vinyl monomers, vinyl cyanide monomers, (meth)acrylate ester monomers and carboxyl group-containing vinyl monomers (Claim 12);

a flame retardant

which comprises the polyorganosiloxane-containing graft copolymer according to Claim 9 (Claim 13);

a resin composition excellent in impact resistance and flame retardancy

which comprises a thermoplastic resin and the flame retardant according to Claim 13 as incorporated therein in an amount of 0.1 to 30 parts by weight per 100 parts by weight of the thermoplastic resin (Claim 14);

[0017] The resin composition according to Claim 14, wherein the thermoplastic resin is a polycarbonate resin (Claim 15).

DETAILED DISCLOSURE OF THE INVENTION

First Aspect of the Invention

[0018] The polyorganosiloxane-containing graft copolymer according to the first aspect of the invention is one obtainable by polymerizing 0.5 to 10 parts of a vinyl monomer (B) (hereinafter also referred to as "vinyl monomer B"). which comprises 100 to 50% of a polyfunctional monomer (b-1) (hereinafter also referred to as "polyfunctional monomer (b-1) ") containing two or more polymerizable unsaturated bonds and 0 to 50% of another copolymerizable monomer (b-2) (hereinafter also referred to as "copolymerizable monomer (b-2)"), in the presence of 40 to 90 parts (as solid content) of polyorganosiloxane particles (A1), followed by further polymerization of 5 to 50 parts of a vinyl monomer

(C), with the sum of (A1), (B) and (C) amounting to 100 parts. [0019] The polyorganosiloxane particles (A1) preferably have a volume average particle diameter of not less than $0.008\,\mu m$, more preferably not less than $0.01\,\mu m$, as determined by light scattering method or electron microscopic observation. On the other hand, it is preferably not larger than 0.6 μm , more preferably not larger than 0.2 μm , most preferably not larger than 0.15 μ m. It tends toward difficulty to obtain particles smaller in volume average particle

diameter than 0.008 μm and, when such diameter exceeds 0.6 μm , the flame retardancy tends to deteriorate. [0020] The coefficient of variation in the particle diameter distribution (100 x standard deviation/volume average particle diameter) (%) of the polyorganosiloxane particles is desirably controlled so that it may amount preferably to 10 to 100%, more preferably to 20 to 60%, since then the moldings produced from the resin composition containing the graft copolymer according to the first aspect of the invention can have a good surface appearance.

[0021] From the flame retardancy and impact resistance viewpoint, the polyorganosiloxane particles (A1) preferably has a toluene-insoluble matter content (as determined by immersing 0.5 g of the particles in 80 ml of toluene at 23 °C

for 24 hours) of not more than 95%, more preferably not more than 50%, most preferably not more than 20%. [0022] In the first aspect of the invention, the polyorganosiloxane particles (A1) include, within the meaning thereof, not only particles made of a polyorganosiloxane(s) alone but also modified polyorganosiloxane particles containing not more than 5% of another or other (co)polymers. Thus, the polyorganosiloxane particles (A1) may contain not more than 5% of polybutyl acrylate, a butyl acrylate-styrene copolymer and/or the like therein.

[0023] As specific examples of the polyorganosiloxane particles (A1), there may be mentioned polydimethylsiloxane particles, polymethylphenylsiloxane particles, and dimethylsiloxane-diphenylsiloxane copolymer particles. These particle species may be used singly or two or more of them may be used in combination.

[0024] The polyorganosiloxane particles (A1) can be obtained, for example, by polymenzing (1) an organosiloxane, (2) a bifunctional silane compound, (3) an organosiloxane and a bifunctional silane compound, (4) an organosiloxane and a vinylic polymerizable group-containing silane compound, (5) a bifunctional silane compound and a vinylic polymerizable group-containing silane compound or (6) an organosiloxane, a bifunctional silane compound and a vinylic polymerizable group-containing silane compound, and the like, and optionally further with an at least trifunctional silane compound (s). The "bifunctional silane compound" is a silane compound having a total number of two of a hydroxyl group (s) and/or a hydrolyzable group(s) each bound to a silicon atom. The "at least trifunctional silane compound" means a silane compound having a total number of at least three of a hydroxyl group(s) and/or a hydrolyzable group

(s) each bound to a silicon atom. [0025] The above-mentioned organosiloxane and bifunctional silane compound are components constituting the main skeleton of the polyorganosiloxane chain. As specific examples of the organosiloxane, there may be mentioned, among others, hexamethylcyclotrisiloxane (D3), octamethylcyclotetrasiloxane (D4), decamethylcyclotrisiloxane

(D5), dodecamethylcyclohexasiloxane (D6), tetradecamethylcycloheptasiloxane (D7), and hexadecamethylcyclohexasiloxane clooctasiloxane (D8). As specific examples of the bifunctional silane compound, there may be mentioned diethoxydimethylsilane, dimethoxydimethylsilane, diphenyldimethoxysilane, diphenyldiethoxysilane, 3-chloropropylmethyldimethoxysilane, 3-glycidoxypropylmethyldimethoxysilane, heptadecafluorodecylmethyldimethoxysilane, trifluoropro-

- Pylmethyldimethoxysilane, oxadecylmethyldimethoxysilane, and the like. From the viewpoint of economical efficacy and flame retardancy, silane compounds or mixtures thereof comprising 70 to 100%, preferably 80 to 100%, of D4 or a mixture of D3 to D6 and, as the rest, to 0 30%, preferably 0 to 20% of diphenyldimethoxysilane, dishbased as the preferably of the preferably 0 to 20% of diphenyldimethoxysilane, dishbased as the preferably of the p
- diphenyldiethoxysilane and of the like are preferred among others.

 [1026] The above-mentioned virilic polymerizable group-containing silane compound is component for introducing a virylic polymerizable or polymerizable group-containing silane compound and/or the above-mentioned virilic polymerizable group into side chairs and/or a terminus or termini of the copolymer by copolymerizable or compound and/or the the above-mentioned organositoxens. Educational silane compound, at least trifunctional silane compound and/or the like, and survivily in colomositic polymerizable group serves as an active site for grafting in chemical binding with a viryl (co) like, and survivily incomer (6) er viryl monomer (C) mentioned later herein. Furthermore, it is a compoun polymer formed from the virylmonomer (8) or viryl monomer (C) mentioned later herein. Furthermore, it is a compound to group and the virylmonomer (8) or viryl monomer (7) mentioned later herein. Capable of forming a crosslink between such active sites of grafting in the manner of radical reaction induced by a radical polymerization initiator may be the same one as can be used in the graft polymerization to the mentioned later herein. Even when crosslinking is caused by the radical reaction, grafting is still possible since such sites partly remain as active sites.
- as clive sites for grating.

 [0027] As specific examples of the above-mentioned virylic polymerizable group-containing silane compound, there
 [0027] As specific examples of the above-mentioned virylic polymerizable group-containing silane compound, among others, mentioned, among others, mentioned, polymerizable group-containing silane, prestancy-loyloxypropylicithoxysilane, propriate and like (methi-port/polyxy group-containing silane comments of the propriate p
- pounds, p-vinylphenyldimethoxymethylsilane, p-vinylphenyltimethoxysilane ara like vinylprenny group-containing siane compounds, vinylmethyldimethoxysilane, vinyltrimethoxysilane, vinyltrimethoxysilane and like vinyl group-containing silane compounds, mercaptopropyltrimethoxymethylsilane and like mercaptogroup-containing silane compounds, mercaptopropyltrimethoxymethylsilane and like mercaptogroup-containing silane compounds, winyl group-containing silane compounds are preferred from the economical viewpoint.
- [0028] In cases where the above-mentioned vinytic polymerizable group-containing slane compound is of the trialkoxysilane type, it serves also as the at least triunctional silane compound mentioned below.
 [0029] The serves also as the at least triunctional silane for introduction a consolinked structure into
- [1022] The at least trifunctional silane compound is used as a component for introducing a crosslinked structure into the polyorganosioxane and providing the same with rubber elasticity as a result of copolymerization thereof with the above-mentioned organosioxane, brifunctional silane compound and/or virigit polymerizable group-containing silane compound, among others, namely as a crosslinking agent for the polyorganosiloxane. As specific examples, there may be mentioned tetraethoxysilane, methyltriethoxysilane, entryltriethoxysilane, entryltriethoxysilane, organization organization of the polyorganosiloxane organization organization of the polyorganosiloxane organization organization
- ysitane are preferably used in view of the high efficiency of crossfinking attainable therewith.

 [0030] In the polymerization thereof, the above-mentioned organositoxane, bifunctional silane compound, vinylic polymerization thereof, the above-mentioned organositoxane compound are generally used in proportions such that the organositoxane and/or bifunctional silane compound (the ratio between the organositoxane and/or bifunctional silane compound (the ratio between the organositoxane and bifunctional silane compound generally being 1000 to 0/100 by weight, preferably 10.00 to 7030 by weight) amounts to 50 to 99.9%, referably 0 to 99%. The vinylic polymerizable group-containing silane compound to 0 to 50%, preferably 0 to 39%, and the at least trifunctional silane compound do not simultaneously polymerizable group-containing silane compound and at least trifunctional silane compound to 0 to 50%, preferably 0 to 39%, and the at least trifunctional silane compound do not simultaneously amount to 0%, and at least one of them is \$\text{Preferably used in an amount of not less than 0.1%.}
- [0031] Whenthe proportion of the above-mentioned organos lovane and/or bifunctional silane compound is too small, the resin composition is existent to be some brittle. When the the resin composition is existent to be composition is existent to be composition in excessive, the amount of the virylic polymerizable group-containing silane compound and/or at least trifunctional silane compound becomes excessively small and the effects of using these tend to be hardly produced. Furthermore, when the proportion of the above-mentioned virylic polymerizable group-containing silane compound and/or at least trifunctional silane compound is too small, the flame retardancy-causing effect becomes unsatisfactory and, when it is excessive, the resin compositions obtained by incorporating the resulting copolymer therein tend to become british.
 - become brittle. [032] The above-mentioned polyorganositoxane particles (A1) are preferably produced, for example, by emulsion [032] The above-mentioned polyorganositoxane particles (A1) are preferably produced, for example, by emulsion polymerization of the polyorganositoxane-forming components, such as the organositoxane, bifunctional silane compound organization of the polyorganositoxane-forming silane compound organization of the polyorganositoxane-forming silane compound as added accompound as add
- compound as added according to need.

 [0033] The above-mentioned emulsion polymerization may be carried out by emulsifying and dispersing the above polyorganosioxene-forming components in water in the presence of an emulsifier by means of mechanical shearing and making the system acidic. When in this case, emulsion droplets not smaller than several micrometers are prepared and making the system acidic. When in this case, emulsion droplets not smaller than several micrometers are prepared by means of mechanical shearing, the voluntaries average particle diameter of the polyorganosiloxane particles (A1) ob-

tained after polymerization can be controlled within the range of 0.02 to 0.6 µm by varying the amount of the emulsifier used. As for the coefficient of variation in particle diameter distribution (100 x standard deviation/volume average particle diff.

ticle diameter) (%), a value of 20 to 70% can be obtained. [0034] For producing polyorganosiloxane particles not larger than 0.1 µm with a narrow particle diameter distribution, the polymerization is preferably carried out in a multistage manner. For example, a 1 to 20% portion of an emulsion composed of emulsion droplets not smaller than several micrometers as obtained by emulsification, under mechanical shearing, of the above-mentioned polyorganosiloxane-forming components, water and the emulsifier is subjected in advance to emulsion polymerization under acidic conditions, the remaining portion of the emulsion is added and subjected to polymenzation in the presence of the polyorganosiloxane particles obtained, which serve as seeds. It is possible to control, by adjusting the emulsifier amount, in a manner such that the thus-obtained polyorganosiloxane particles may have a volume average particle diameter of 0.02 to 0.1 μm with a coefficient of variation in particle diameter distribution of 10 to 60%. According to a more preferred procedure for the multistage polymenzation, the multistage polymerization is carried out in the same manner using a vinyl (co)polymer obtained by ordinary emulsion polymerization of a vinyl monomer(s) (e.g. styrene, butyl acrylate, methyl methacrylate), which is(are) used on the occasion of graft polymerization mentioned later herein, in lieu of the polyorganosiloxane particles serving as seeds, whereby it is possible to control, by adjusting the emulsifier amount, in a manner such that the polyorganosiloxane (modified polyorganosiloxane) particles obtained may have a volume average particle diameter of 0.008 to 0.1 µm with a coefficient of variation in particle diameter distribution of 10 to 50%. The above-mentioned emulsion droplets not

smaller than several micrometers can be prepared by using a high-speed stirrer, for example a Homorrivation [0033] In the above-mentioned emulsion polymerization, an emulsifier which will not lose its emulsifying ability under locality of the state of

serve as polymerization catalysts for the polyorganosiloxane-forming components.

[038] The acidic condition can be obtained by adding, to the system, an inorganic acid, such as sulfuric acid or hydrochloric acid, or an organic acid, such as an alkylbenzesulfonic acid, an alkylsulfonic acid or trifluoroacetic acid. Hydrochloric acid, or an organic acid, such as an alkylbenzesulfonic acid, an alkylsulfonic acid or trifluoroacetic acid. For avoiding corrosion of the production equipment and for attaining an adequate rate of polymerization, the pH is

preferably adjusted to 1 to 3, more preferably to 1.0 to 2.5.

[0037] For attaining an adequate rate of polymerization, the heating for the polymerization is carried out preferably at 60 to 120°C, more preferably 70 to 100°C.

[0038] Under such an acidic condition, the Si-O-Si bonds forming the polyorganosiloxane skeleton are in a state of sequilibrium between cleavage and formation, and this equilibrium varies depending on the temperature. Therefore, for equilibrium between cleavage and formation, and this equilibrium varies depending on the temperature lowers, the stabilizing the polyorganosiloxane chain, the system is preferably neutralized with an aqueous solution of an alkali, such as sodium hydroxide, potassium hydroxide or sodium carbonate. Furthermore, as the temperature lowers, the above equilibrium shifts to the formation side, facilitating the formation of products with a high-molecular weight and a high-degree of crosslinking, so that, for obtaining polymer having high-molecular weight or high degree of crosslinking, so that, for obtaining polymer having high-molecular weight or high degree of crosslinking, and the system after carrying out the polymerization of the polyorganosiloxane-forming components at 60°C or above is

preferably cooled to nom temperature or below and, after about 5 to 100 hours of standing, it is neutralized.

[0039] The thus-obtained polyorganosiloxane particles (A1), when formed, for example, by polymertzation of the organosiloxane and/or bifunctional silane compound, with or without further addition of the virylic polymertzable group-containing silane compound, occur as a virylic polymertzable group-containing polymer generally resulting from random copolymerization. In cases where the at least trifunctional silane compound is used in the copolymerization, the armound of the control of the con

tween vinylic polymerizable groups. with part of the vinylic polymerizable groups remaining unreacted.

[0040] Graft polymerization of the vinyl monomer (6) and vinyl monomer (C) onto the polyorganosiloxane particles

(A1) gives the polyorganosiloxane-contraining graft copolymer. In the polymerization of the vinyl monomer (8) and vinyl

(A1) gives the polyorganosiloxane-contraining graft copolymer. In the polymerization of the vinyl monomer (6) and vinyl

alled free polymer molecules resulting from polymerization of the branch components (herein polymers of the vinyl

monomer (6) and vinyl monomer (C)) alone without their grafting onto the stem component (herein the polyorganosil
monomer (8) and vinyl monomer (C)) alone without their grafting onto the stem component (herein the polyorganosil
monomer (8). The product is thus obtained as a mixture of the graft copolymer molecules.

In accordance with the first appeal of the vention, these both species are collectively referred to as the graft copolymer is not the vinyl monomer (B) onto the

polyorganosiloxane particles (A1) and graftling of the vinyl monomer (C) not only onto the polyorganosiloxane particles (A1) and graftling of the vinyl monomer (C) not only onto the polyorganosiloxane particles (A1) and graftling of the vinyl monomer (C).

(A1) but also onto the polymer molecules formed by the vinyl monomer (B), hence the free polymer content is low. [0042] The graft copolymer preferably has an acetone-insoluble matter content (as determined by immersing 1 g of the graft copolymer preferably not less than 85%, more preferably not less than 85% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 85% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 85% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 85% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 85% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 85% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80%, more preferably not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80% in view of copolymer in 80 ml acetone at 23 °C for 48 hours) of not less than 80% in view of copolymer in 80 °C for 48 hours).

in view of good flame retardant effects obtainable in such a case. [0043] The above-mentioned vinyl monomer (B) is used for enhancing the flame retardant and impact resistanceimproving effects. It comprises 100 to 50%, preferably 100 to 80%, more preferably 100 to 90%, of a polyfunctional monomer (b-1) containing two or more polymerizable unsaturated bonds and 0 to 50%, preferably 0 to 20%, more preferably 0 to 10%, of another copolymerizable monomer (b-2). When the proportion of the polyfunctional monomer (b-1) is too small, or when the copolymerizable monomer (b-2) is excessive, the graft copolymer finally obtained tends to become

to become less effective in improving the impact resistance [0044] The polyfunctional monomer (0-1) is a compound containing two or more polymerizable unsaturated bonds in the molecule. As specific examples thereof, there may be mentioned allyl methacrylate, triallyl cyanurate, triallyl isocyanurate, diality phthalate, ethylene glycol dimethacrylate, 1,3-butylene glycol dimethacrylate, and divinylbenzene. These may be used singly or two or more of them may be used in combination. Among them, the use of allyl methacrylate is

ylate, in particular, is preferred from the economical and efficacy viewpoint. [0045] As specific examples of the copolymerizable monomer (b-2), there may be mentioned, among others, aromatic whole matic vinyl monomers such as styrene, a methylstyrene, paramethylstyrene and parabutylstyrene, vinyl cyanide monomers such as styrene, a methylstyrene, paramethylstyrene and parabutylstyrene, vinyl cyanide monomers such omers such as acrylonitrile and methacrylonitrile, (meth)acrylate ester monomers such as methyl acrylate, ethyl acrylate ster monomers such as acrylonitrile and methacrylonitrile. Ylate, propyl acrylate, butyl acrylate, 2-ethylinexyl acrylate, glycidyl acrylate, hydroxyethyl acrylate, hydroxyethyl acrylate, butyl acrylate, 2-ethylinexyl acrylate, 1-ethylinexyl acrylate, 1-ethylinexyl acrylate, 1-ethylinexyl acrylate, 2-ethylinexyl acrylate, 1-ethylinexyl acrylate, 1-ethylinexyl acrylate, 2-ethylinexyl acrylate, 2-ethylinexyl ylate, methyl methacrylate, butyl acrylate, 2-eurymeen, methyl methacrylate, lauryl methacrylate, glycidyl methacrylate and hy-droxyethyl methacrylate, ethyl methacrylate, butyl methacrylate, butyl methacrylate, size of fundamental methacrylate and fundamental methacrylate. droxyethyl methacrylate, ethyl methacrylate, one carboxyl group-containing vinyl monomers such as itaconic acid, (meth)acrylic acid, fumaric acid and maleic acid. These may be used singly or two or more of them may be used in combination.

[0046] The above-mentioned vinyl monomer (C) is a component to be used for obtaining the polyorganosilloxanecontaining graft copolymer. It is also a component to be used for securing compatibility between the graft copolymer and a thermoplastic resin to thereby disperse the graft copolymer in the thermoplastic resin uniformly for the improvements in flame retardancy and impact resistance by incorporating the graft copolymer in the thermoplastic resin. There-fore, the visit recovery and impact resistance by incorporating the graft copolymer of the visit recovery resistance and the property resistance fore, the viryl monomer (C) is preferably selected so that a polymer of the viryl monomer may preferably have a Solubility or loss than 9 17 (calicm³)1/2, still more preferably not loss than 9 17 (calicm³)1/2. solubility parameter of not less than 9.15 (call/cm³)1/2, more preferably not less than 9.17 (call/cm³)1/2, still more preferably not less than 9.15 (call/cm³)1/2, it is call-and to that the solubility parameter is question may be erably not less than 9.10 (Cal/cm3)1/2. Also preferably, it is selected so that the solubility parameter in question may be not less than 9.20 (Cal/cm3)1/2. Also preferably, it is selected so that the solubility parameter in question may be not less than 10.10 (cal/cm3)1/2. etill more preferably not more than not less than 10.10 (cal/cm3)1/2. not more than 10.15 (cal/cm³)^{1/2}. Also pretrainly not more than 10.10 (cal/cm³)^{1/2}, still more preferably not more than 10.10 (cal/cm³)^{1/2}, more preferably not more than 10.10 (cal/cm³)^{1/2}, and the flower called 10.05 (cal(cm³) 1/2, more preterant, 10.05 (cal(cm³) 1/2. When the solubility parameter is outside the above range, the flame retardancy tends to decrease. [0047] The solubility parameter values are calculated using the small group parameters according to the group contribution multiplication and the solubility parameter values are calculated using the small group parameters according to the group contribution multiplication and the solubility parameter values are calculated using the small group parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution multiplication and the solubility parameters according to the group contribution and the group contribution according to the group contribution and the group contribution according to the group contribution and the group contribution according to the group contribution and the group contribution according to th Inbution method described in "Polymer Handbook", 4th edition, published by John Wiley & Sons, Inc., 1999, Section Will, pages 699 of the Control of the Cont VII, pages 682-685. For example, the value for poly (methyl methacrylate) (regarding the repeating unit molecular weight as 100 method is 9.75 freal/cm3\1121. for poly/butyl acrylate) (regarding the weight as 100 g/mole, and the density as 1.19 g/cm³) is 9.25 [(cal/cm³)^{1/2}], for poly(butyl acrylate) (regarding the repeating unit molecular weight as 1.18 g/mole, and the density as 1.06 g/cm³) 8.97 [(cal/cm³) 17], for poly(buty) meth-acrylate) (or a significant weight as 128 g/mole, and the density as 1.06 g/cm³) 9.47 [(cal/cm³) 14.06 g/cm³) 9.47 [(cal/cm³) 17], for poly(buty) methacrylate) (regarding the repeating unit molecular weight as 142 g/mole, and the density as 1.06 g/cm³) 9.47 ((cal/ cm³)^{1/2}, for expectating unit molecular weight as 104 and the density as 1.05 g/cm³) 9.03 cm³/12], for polystyrene (regarding the repeating unit molecular weight as 104, and the density as 1.05 g/cm³) 9.03 [[Cal/cm³/12]]. [(cal/cm⁵)¹/2, and for polyacrylonitrile (regarding the repeating unit molecular weight as 53, and the density as 1.18 g/ cm³ 1.271 (cm³ 1.271) (cm cm³) 12.71 [(cal/cm³)^{1/2}]. Used as the density values for the respective polymers were those described in Ullman's Encyclopedia. Encyclopedia of Industrial Chemistry, published by VCH, 1992, volume A21, page 169. As for the solubility parameter for each post of ea So of each copolymer, the value for the main component was employed when the copolymer weight fraction is less than 5% and when the additivity rule based on the weight than 5% and, when that weight fraction is not less than 5%, it was supposed that the additivity rule based on the weight fractions are held. fractions can hold good. Thus, the solubility parameter & can be calculated from the solubility parameters on frespective homopolymers of m vinyl monomer species constituting the copolymer in question and weight fraction Wn of that polymer according that polymer according to the following equation (1):

$$\delta_{c} = \sum_{n=1}^{n=m} \delta_{n} W_{n} / \sum_{n=1}^{n=m} \delta_{n}$$
(1)

nitrile can be found to be 9.5 [(cal/cm³)^{1/2}] by using the solubility parameter 9.03 ((cal/cm³)^{1/2}] of polystyrene and the solubility narameter 9.03 ((cal/cm³)^{1/2}) of polystyrene 9.03 ((cal/cm³)^{1/2}) by using the solubility narameter 9.03 ((cal/cm³)^{1/2}) of polystyrene 9.03 (solubility parameter 12.71 [(cal/cm³) 1/2] by acrylonitrile and using the equation (1).

[0049] For the solubility parameter &s of a viryl polymer obtained by carrying out polymerization in two or more stages while varying the vinyl monomer species in each stage, it was supposed that the additivity rule can hold good with respect to the values obtained by dividing the weights of the vinyl polymers obtained in the respective stages by the whole weight of the vinyl polymer finally obtained, namely the weight fractions. Thus, the value &s of the polymer polymerized in q stages can be calculated from the solubility parameter δi of the polymer obtained in each stage and the weight fraction Wi of that polymer according to the following equation (2):

$$\delta_{s} = \sum_{i=1}^{i=q} \delta_{i} W_{i} / \sum_{i=1}^{i=q} W_{i}$$
(2)

[0050] Thus, for example, when polymerization was carried out in two stages and, in stage 1, 50 parts of a copolymer composed of 75% of styrene and 25% of acrylonitrile was obtained and, in stage 2, 50 parts of a polymer of methyl methacrylate was obtained, the solubility parameter of the polymer obtained by such two-stage polymenzation can be calculated as 9.60 [(cal/cm³)1/2] by using the solubility parameter value 9.95 [(cal/cm³)1/2] for the 75% styrene-25% acrylonitrile copolymer and the solubility parameter value 9.25 [(cal/cm³)1/2] for poly(methyl methacrylate) and using

[0051] The above-mentioned vinyl monomer (C) includes the same ones as mentioned hereinabove as the other copolymerizable monomer (b-2) referring to the vinyl monomer (B). These may be used singly or two or more of them may be used in combination.

- [0052] The polyorganosiloxane-containing graft copolymer according to the first aspect of the invention can be obtained by polymerizing 0. 5 to 10 parts (preferably not less than 1 part, more preferably not less than 2 parts, but preferably not more than 5 parts, more preferably not more than 4 parts) of the vinyl monomer (B) in the presence of 40 to 90 parts (as solid content) (preferably not less than 60 parts, but preferably not more than 80 parts, more preferably not more than 75 parts) of the above-mentioned polyorganosiloxane particles (A1) and further polymerizing 5 to 50 parts (preferably not less than 15 parts, more preferably not less than 21 parts, but preferably not more than 39 parts, more preferably not more than 38 parts) of the vinyl monomer (C) so that the total amount may become 100 parts. When the amount of the polyorganosiloxane particles (A1) is too small or too large, the flame retardant effect tends to decrease in either case. When the amount of the vinyl monomer (B) is too small, the flame retardant and impact resistance improving effects tend to decrease and, when it is excessive, the impact resistance improving effect tends to decrease. When the vinyl monomer (C) is too small or too large, the flame retardant effect tends to lower in either case. [0053] In carrying out above graft polymerization, the technique of ordinary seed emulsion polymerization can be applied. Thus, the above-mentioned vinyl monomer (B) and vinyl monomer (C) can be subjected to radical polymeration to the subject of the sub zation in the presence of polyorganosiloxane particles (A1). In the radical polymerization in question, the polyorganosiloxane nosiloxane particles (A1) are preferably in a latex form. Namely, it is preferred that the radical polymerization should be carried out in latex of the polyorganosiloxane particles (A1). The vinyl monomer (B) and vinyl monomer (C) each
- may be polymerized in one stage or in two or more stages. [0054] The above radical polymerization can be carried out by the method comprising thermally decomposing a radical polymerization initiator to thereby cause the reaction to proceed, or by the method comprising allowing the reaction to proceed in a redox system using a reducing system, for instance, without any particular restriction.
- [0055] As specific examples of the radical polymerization initiator, there may be mentioned organic peroxides such as cumene hydroperoxide, tert-butyl hydroperoxide, benzoyl peroxide, tert-butyl peroxylsopropyl carbonate, di-tert-butyl butyl peroxide, tert-butyl nydroyl peroxide, succinic acid peroxide, cyclohexanone peroxide, and acetylacetone peroxide, inorganic peroxides such as potassium persulfate and ammonium persulfate, and azo compounds such as 2.2 azobisisobutyronitrile and 2.2 azobis-2.4-dimethylvaleronitrile, among others. Among these, organic peroxides and inorganic peroxides are particularly preferred because of their high reactivity.
- [0056] As the reducing agent to be used in the above-mentioned redox system, there may be mentioned such mixture as iron(II) sulfate/glucose/sodium pyrophosphate, iron(II) sulfate/dextrose/sodium pyrophosphate, and iron(II) sulfate/ sodium formaldehyde sulfoxylate/ethylenediamine acetate, for instance.
- [0057] The above-mentioned radical polymerization initiator is used generally in an amount of 0.005 to 20 parts, preferably 0.01 to 10 parts, most preferably 0.04 to 5 parts, per 100 parts of the sum total of the vinyl monomer (B) and/or vinyl monomer (C) used or, in the case of multi-stage polymerization, per 100 parts of the monomer(s) used in each stage. In multi-stage polymerization, the radical polymerization initiator and the amount thereof each may be the same or different in each stage. When the amount of the radical polymerization initiator is too small, the rate of reaction is low and, accordingly, the production efficiency tends to worsen. When it is excessive, the generation of heat during

reaction tends to become intensive, making it difficult to produce the desired graft copolymer.

[0058] A chain transfer agent may also be used where necessary in carrying out the radical polymerization. The chain transfer agent may be any of those generally used in emulsion polymerization, without any particular limitation. [0059] As specific examples of the chain transfer agent, there may be mentioned tert-dodecylmercaptan, n-octylmercaptan, activatedecylmercaptan, and n-hexylmercaptan, among others.

ercaptan, n-tetradecyimercaptan, arid n-texymercaptan, ariding useries.

[0660] Although it is an optional component, the chair transfer agent is used preferably in an amount of 0.01 to 5 parts per 100 parts of the sum of the vinyl monomer (B) and/or vinyl monomer (C) employed. In multi-stage polymercation, it is used preferably in an amount of 0.01 to 5 parts per 100 parts of the monomer (s) used in each stage. In multi-stage polymercation, the chair transfer agents and the addition levels thereof in the respective steps may be the same or different. When the amount of the chair transfer agent is smaller than 0.01 parts, no significant effect can be obtained and, when it exceeds 5 parts, the rate of polymerization slows down, hence the production efficiency tends

[0061] Generally, it is preferred that the reaction temperature be 30 to 120°C.

5

[0062] When, in the above polymerization, the polyorganosiloxane particles (A1) contain vinylic polymerization groups, the vinyl monomer (B), on the occasion of polymerization thereof by means of the radical polymerization initiator, reacts with the vinylic polymerizable groups of the polyorganosiloxane particles (A1) to form grafts. When the polyorganosiloxane particles (A1) have no vinylic polymerizable group, a specific radical initiator, for example tert-butyl granosiloxane particles (A1) the vinyl monomer (B) is polymerizad by the resulting radicals to form grafts. Furthermore, when the vinyl monomer (B) is polymerizad by the resulting radicals to form grafts. Furthermore, when the vinyl monomer (C) is polymerized by a radical polymerization initiator, it reacts not only with the polyorganosiloxane particles (A1), like the vinyl monomer (B), but also with the unsaturated bonds occurring in the polymer molecules formed by the vinyl monomer (B) to give grafts resulting from the vinyl monomer (C).

[0663] The graft copolymer produced by emulsion polymerization may be separated from the latex for use thereof, or may be used in the latex form. The method of recovering the polymer from the latex may be any of conventional methods. For example, mention may be made of the method comprising adding a metal salt, such as calcium chloride, magnesium chloride or magnesium cultate, to the latex to cause the latex to coagulate, followed by separation, washing with water, dehydrating and drying. The spary drying method may also be used.

[0064] The thus-obtained graft copolymer is incorporated in various thermoplastic resins to give flame-retardant resin compositions excellent in flame retardancy and impact resistance.

composuons exceient in itame retrialative and impact resistance.

[0055] Preferred as the thermoplastic resistance are polycarbonate resins whose polycarbonate content is not tess than 50%, more preferably not less than 70%, since good flame retardancy can be obtained with them. Specific examples of the thermoplastic resists, which are preferred from the economical viewpoint and in view of good balance between flame retardancy and impact resistance, are polycarbonates (in particular aromatic polycarbonates), polycarbonated polycarbonates in such as polycarbonated polycarbonated polycarbonated polycarbonated polycarbonated bend resis and polycarbonated/polybutylene terephthalate blend resins, polycarbonate/polybutylene terephthalate blend resins, polycarbonate/pultidiene-styrene copolymer (HIPS resin) blend resins, polycarbonate/acrylonitrile-styrene copolymer (Bals resin) blend resins, polycarbonate/acrylonitrile-styrene copolymer blend resins, polycarbonate/styrene copolymer (LIPS resin) blend resins. Mixtures of two or more blend resins and polycarbonate/acrylonitrile-N-phenylmaleimide copolymer blend resins, and polycarbonate/acrylonitrile-A-phenylmaleimide copolymer blend resins, and polycarbonate/acrylonitri

45 (2007) The powder-form flame retardant comprising the polyorganosiloxane-containing graft copolymer separated from the latex can be admixed with the thermoplastic resins by mixing using a Henschel mixer or ribbon blender, for instance, followed by melting and kneading using a roll, extruder or kneader, for instance.

[0068] On that occasion, one or more of additives in general use, namely antioxidants, dripping-preventing agents, polymer processing auxiliaries, flame retardants, impact resistance improver, plasticizers, lubricants, ultraviolet absorbers, pigments, glass fibers, fillers, polymer lubricants and so forth, may be incorporated in the resin compositions. [0069] As specific examples of the antioxidants, there may be mentioned, among others, phenolic antioxidants such as trisfN-(3,5-di-tert-butyl-4-hydroxypheny)[propionyloxymethyl]methane (e.g. IRGANOX 1010, product of ASAH IDENKA), lett-alsiyl-4-di-driv-butyl-4-hydroxypheny)[propionyloxymethyl]methane (e.g. IRGANOX 1010, product of Clba Special-ty Chemicals), butylidene-1,1-bis(2-methyl-4-hydroxy-5-tert-butylpheny)[butane (e.g. Yoshinox 39a), product of ASAH IDENKA), and 1,1-3-tris(2-methyl-4-hydroxy-5-tert-butylpheny)[butane (e.g. Yoshinox 39a), product of Yoshinomi Fine Chemicals), phosphorus-containing antioxidants such as bis(2,6-di-tert-butyl-4-methylpheny)) pentaerythritol phosphite (e.g. ADEKA STAB PLP-36, product of ASAH IDENKA) and 2,2-methylenebis(4,6-di-tert-butyl-heny)) octyl phosphite (e.g. ADEKA STAB 212-product of ASAH IDENKA) and 2,2-methylenebis(4,6-di-tert-butyl-heny)) octyl phosphite (e.g. ADEKA STAB 212-product of ASAH IDENKA) and 2,2-methylenebis(4,6-di-tert-butyl-heny)) octyl phosphite (e.g. ADEKA STAB 212-product of ASAH IDENKA) and 2,2-methylenebis(4,6-di-tert-butyl-heny)) octyl phosphite (e.g. ADEKA

STAB HP-10, product of ASAHI DENKA), and sulfur-containing antioxidants such as dilauryl 3,3'-thio-dipropionate (Yoshinox DLTP, product of Yoshitom Fine Chemicals) and dimyristyl 3,3'-thio-dipropionate (Yoshinox DMTP, product of Yoshitomi Fine Chemicals), Among these, phosphorus-containing antioxidants are particularly preferred since they

- provide improved fiame retardancy.

 [10070] As examples of the dripping-preventing agents, which are preferred in view of their high dripping-preventing agents, which are preferred in view of their high dripping-preventing effect, there may be mentioned fluorinated polydefin resins such as polymonofluoroethylene, polydirfluoroethylene, polyterafluoroethylene and tetrafluoroethylene/exafluoroethylene polyterafluoroethylene and tetrafluoroethylene and tetrafluoroethylene are fluoridely resins.
- (0071) As specific examples of the polymer processing auxiliaries, there may be mentioned, among others, methyl methacrylate butyl acrylate copolymers and like methacrylate (co)polymers.
 - [0072] As specific examples of the impact resistance improvers, there may be mentioned, among others, butadiene rubber type impact resistance improvers, and rubber type impact resistance improvers, and butyl acrylate rubber stance improvers (MBS resins), butyl acrylate rubber stance improvers. Butyl acrylate rubber/silicone rubber and like composite rubber type impact resistance improvers.
 - [0073] One or more of other fame relatdants may also be used in combination. As specific examples of the fame relatdants to be used in combination, which are preferred from the fact that they are hadopen-free and phosphorus free, there may be mentioned, among others, slicone compounds such as aromatic group-containing polyorganositoxanes, triazine compounds such as varunfue acid and melamine cyanurate, and boron compounds such as brond loxanes, triazine compounds such as varunfue acid and melamine cyanurate, and boron compounds such as they hope produced to the properties of the product of the product of the product of the polyorganositoxanes, triazine compounds such as triphable, concluded and zinc borate. The combined use with a phosphorus is also possible. In this case, the use of the polyorganositoxane-containing grather esters or stabilized red phosphorus is also possible. In this case, the use of the polyorganositoxane-containing a phosphorus containing and prosphorus containing and prosphorus containing and prosphorus desired productions.
- reduced thereby.

 [0074] From the effect-cost balance viewpoint, the level of addition of these additives is preferably 0.1 to 20 parts,
 more preferably 0.2 to 10 parts, and most preferably 0.3 to 5 parts, per 100 parts of each thermoplastic resin.
- with preterably 0.2 to 10 parts, and most preterably 0.3 to 40 parts, and most preterably 0.3 to 10 parts, and most preterably 0.3 to 10 parts, and most preterable of the molding and so furth.
- [0076] The fields of application of the moldings obtained from the flame retardant resin composition according to the first aspect of the invention are not particularly restricted but include those fields where flame retardancy is required, for example housings and chassis parts of various OA/findmation/household electric-felectronic appliances such as desktop computers, notations computers, tower type computers, server computers, printers, copiers, fax machines, callular phones, PHS phones, televisions and video recorders, various building parts and various automotive parts. [0077] The moldings obtained show excellent impact resistance and flame retardancy.

35 Second aspect of the invention

- [0078] The polyorganosiloxane-containing graft copolymer according to the second aspect of the invention is obtainable by polymerizing 0 to 10 parts of a vinyl monomer (8) comprising 100 to 5% of a polyfunctional monomer (1) containing two or more polymerizable unsaturated bonds and 0 to 50% of another copolymerizable monomer (b. 2). In the presence of 30 to 95 parts of a polyorganosiloxane (A2) in a latex form obtainable by seed polymerization, as a seed polymer, a hydrophilic polymer capable of swelling in the corresponding organosiloxane (such polyorganosiloxane (a2)), followed by further polymerization of 5 to organosiloxane (a2) in the present of a vinyl monomer (c) (the surn of (A2), (8) and (C) being 100 parts).
- [0079] The seed polymer to be used in accordance with the second aspect of invention can be obtained by ordinate of sufficiently restricted. The seed polymer is not finited to such a rubber or but display in the seed polymer such as a butyl acrylate such a rubber component as butyl acrylate rubber or but display or but display or butyl acrylate-styrene copolymer, butyl acrylate-but addiene copolymer, butyl acrylate-butylatione copolymer may also be used without any substantial troub. However, acrylonitrile copolymer or styrene-acrylonitrile copolymer is equired to be capable of sufficiently swelling in the monomer(s) forming a rubber in the next stage the seed polymer is required to be capable of sufficiently swelling in the nonomer(s) forming a rubber in the next stage and have strong hydrophicity so that it may take up water into particles thereof.
- [0080] As a method for improving the hydrophilicity of seed polymer, lowering the glass transition point of the seed polymer may be mentioned. The glass transition point is preferably not higher than 10°C. The glass transition point can be determined, by measuring temperature variation of specific gravity of a polymer, as the temperature at which the specific gravity is drastically useful.
- (9081) For improving the ability of the seed polymer to swell in the organosiloxane, it is first important that the polarity or the like property of the seed polymer is adapted to the organosiloxane. Secondly, it is effective to markedly reduce the molecular weight of the seed polymer by selecting the use of a chain transfer agent, applying a high polymerization temperature and/or use of the initiator in a large amount. A number average molecular weight of the seed polymer is

preferably not more than 10,000, more preferably 7,000. The number average molecular weight may be determined by GPC analysis (relative to polystyrene standard).

[0082] The "organosiloxane", associated with the swelling capacity of the seed polymer, is an organosiloxane which is a monomer component of a polyorganosiloxane (A2). For example, when a polyorganosiloxane (A2) is produced from octamethylcyclotetrasiloxane, the above "organosiloxane" corresponds to octamethylcyclotetrasiloxane.

[0083] The hydrophilicity of the seed polymer can be determined by adding water, in an amount of 20 times (by weight) that of the seed polymer in a dry state, to the dry seed polymer, stirring the mixture at 23 °C for 1 hour, and measuring the extraction rate of the polymer into water. When the value is not less than 1%, the hydrophilicity is sufficient. Preferably, however, the value is not less than 10%, more preferably not less than 50%. The upper limit may

[0084] The swelling capacity of the seed polymer to the organosiloxane can be determined by adding the organosiloxane, in an amount 50 times (by weight) that of the dry seed polymer, to the seed polymer latex, stirring the mixture at 23 °C for 1 hour, and determining the rate of swelling by volume from the ratio between the latex particle diameter after stirring and the latex particle diameter before stirring. A value not less than 1.5 times is sufficient. Preferably, however, it is not less than 3. The upper limit is preferably set at a level not higher than 50 times, more preferably not

[0085] Marked flame retardancy and impact resistance improving effects can be obtained within the above-mentioned

[0086] The volume average particle diameter of the polyorganosiloxane (A2) in a latex form can be determined by light scattering method or electron microscopic observation, and a preferred range is 0.008 to 0.6 µm. More preferably, it is not less than 0.01 μm . As for the upper limit, the diameter is more preferably not greater than 0.2 μm . Those particles smaller than 0.008 μ m in volume average particle diameter tend to become difficult to produce. When the particles are greater than 0.6 µm, the flame retardancy and impact resistance tend to decrease.

[0087] In accordance with the second aspect of the invention, the polyorganosiloxane (A2) includes, within the meaning thereof, not only simple polyorganosiloxanes but also modified polyorganosiloxanes containing not more than 5% of another or other (co)polymers. Thus, the polyorganosiloxane (A2) may contain up to 5% of poly(butyl acrylate) and/ or a butyl acrylate-styrene copolymer, for instance.

[0088] As specific examples of the polyorganosiloxane (A2), there may be mentioned polydimethylsiloxane particles, polymethylphenylsiloxane particles, dimethylsiloxane-diphenylsiloxane copolymer particles, and the like. These may

be used singly or two or more of them may be used in combination. [0089] The above polyorganosiloxane (A2) can be prepared, for example, by polymerizing (1) an organosiloxane, (2) a bifunctional silane compound, (3) an organosiloxane and a bifunctional silane compound, (4) an organosiloxane and a vinylic polymerizable group-containing silane compound, (5) a bifunctional silane compound and a vinylic polymerizable group-containing silane compound, or (6) an organosiloxane, a bifunctional silane compound and a vinylic polymerizable group-containing silane compound, optionally further together with an at least trifunctional silane compound. As these compounds, those specific examples mentioned hereinabove referring to the first aspect of the invention may be mentioned. The proportions thereof are also the same as mentioned hereinabove referring to the first

[0090] The polyorganosiloxane (A2) is preferably produced by subjecting, to emulsion polymerization in the presence of the above-mentioned seed polymer, a polyorganosiloxane-forming composition comprising, for example, the organosiloxane, bifunctional silane compound, vinylic polymerizable group-containing silane compound and/or the like, optionally together with the at least trifunctional silane compound. The method of production is the same as mentioned above referring to the first aspect of the invention.

[0091] The above-mentioned vinyl monomer (B) is used for enhancing the flame retardant and impact resistanceimproving effects. It comprises 100 to 50%, preferably 100 to 80%, of a polyfunctional monomer (b-1) containing two or more polymenzable unsaturated bonds and 0 to 50%, preferably 0 to 20%, of another copolymerizable monomer (b-2). When the proportion of the polyfunctional monomer (b-1) is too small, or when the amount of the copolymenzable monomer (b-2) is excessive, the graft copolymer finally obtained tends to become less effective in improving the impact resistance. As examples of the polyfunctional monomer (b-1) and of the copolymerizable monomer (b-2), there may

be mentioned the same ones as mentioned hereinabove referring to the first aspect of the invention. [0092] The above-mentioned vinyl monomer (C) is a component to be used for obtaining the polyorganosiloxane containing graft copolymer. It is also a component to be used for securing compatibility between the graft copolymer and a thermoplastic resin to thereby disperse the graft copolymer in the thermoplastic resin uniformly for the improvements in flame retardancy and impact resistance by incorporating the graft copolymer in the thermoplastic resin. As specific monomers, there may be mentioned the same ones as mentioned above as the other copolymerizable monomers, there may be mentioned the same ones as mentioned above as the other copolymerizable monomers.

omers (b-2) included in the vinyl monomer (B)-[0093] The polyorganosiloxane-containing graft copolymer according to the second aspect of the invention can be obtained by polymenzing 0 to 10 parts (preferably not less than 1 part, more preferably not less than 2 parts, but

preferably not more than 5 parts, more preferably not more than 4 parts) of the vinyl monomer (B) in the presence of 05 parts, (as solid content) (preferably not less than 60 parts, but preferably not more than 80 parts, more preferably not more than 75 parts) of the above-mentioned obyerganosiloxane (A2) and further polymerizing 5 to 70 parts (preferably not less than 15 parts, more preferably not less than 15 parts, more preferably not less than 21 parts, but preferably not more than 39 parts, more preferably not less than 15 parts, more in 10 parts. When the amount of the polyorganosiloxane (A2) is too small or too large, the flame retardant effect tends to decrease in the amount of the vinyl monomer (B) is too small, the flamer retardant and impact resistance improving effect tends to decrease. When the successive, the impact resistance improving effect tends to decrease. When the successive, the successive preferably monomer (C) is too small or too large, the flame retardant effect tends to lower in either case. Under the form the preferably monomer (C) is too small or too large, the flame retardant effect tends to lower in either case.

first aspect of the invention.

[0995] As the method of recovering the polymer from the graft copolymer latex obtained by emulsion polymerization, [0995]. As the method of recovering the polymer from the graft copolymer latex obtained by emulsion polymerization, there may be mentioned, for example, the method comprising adding a metal salt, such as calcium chloride, magnesium there may be mentioned, for example, the method comprising adding a metal salt, such as calcium chloride, magnesium suifate, to the latex to cause the latex to coagulate, followed by separation, washing with water, chloride or magnesium suifate, to the latex to cause the latex to coagulate, followed by separation, washing with water,

dehydrating and drying. The spray drying method may also be used. [0996] The thus-obtained graft coppymer is incorporated in various thermoplastic resins to give flame-retardant resin compositions excellent in flame retardancy and impact resistance. As examples of the thermoplastic resins, there may

be mentioned the same ones as mentioned hereinabove referring to the first aspect of the invention.

(0937) The level of addition of the above-mentioned polyorganosiloxane-containing graft copolymer to such a ther(0937) The level of addition of the above-mentioned polyorganosiloxane-containing graft copolymer to such a thermoplastic resin is preferably 0.1 to 30 parts per 100 parts of the themcoplastic resin from the good flame retardancy
and impact resistance veworint. More preferably, the addition level is not less than 0.5 parts, still more preferably not

less than 1 part. It is more preferably not more than 15 parts, still more preferably not more than 5 parts. [0099] The powder-form flame retardant comprising the polyorganosiloxane-containing graft copolymer separated from the latex can be admixed with the thermoplastic resins by mixing using a Henschel mixer or ribbon blender, for

instance, followed by melting and kneading using a roll, extruder or kneader, for instance. [Glowed by melting and kneading using a roll, extruder or kneader, for instance. [Glowed by melting and kneading using a roll, extruder or kneader, for instance, improver, plasticizers, lubricans, ultraviolet abpolymer processing auxiliaries, flame retardants, impact resistance improver, plasticizers, lubricans, ultraviolet abpolymer processing auxiliaries, flame retardant and so forth, may be incorporated in the resi compositions, sorbers, pigments, glass fibers, fillers, polymer lubricants and so forth, may be incorporated in the resi compositions, sorbers, pigments, glass fibers, fillers, polymer lubricants and so forth, may be incorporated in the resin composition obtained can be molded by applying those molding methods used in [9100]. The flame retardant resin composition obtained can be molded by applying those molding methods used in

molding conventional thermoplastic resin compositions, namely injection molding, extrusion molding, blow molding, calender molding and so forth.

[1011] The fields of application of the moldings obtained from the flame retardant resin composition according to the ground support of the invention are not particularly restricted but include those fields mentioned hereinabove referring second aspect of the invention are not particularly restricted but include those fields mentioned hereinabove referring

to the first aspect of the invention.

[0102] The moldings obtained show excellent impact resistance and flame retardancy.

BEST MODES FOR CARRYING OUT THE INVENTION

[0103] The following examples illustrate the invention more specifically. They are, however, by no means limitative of the scope of the invention. In the following examples and comparative example, the measurements and tests were carried out as follows.

[Degree of conversion in polymerization]

[0104] The latex was dried in a hot air drying chamber at 120 °C for 1 hour, the remaining solid matter was weighed, and the degree of conversion was calculated according to the formula: 100 x solid matter weight/charged monomer weight (%).

50 [Toluene-insoluble matter content]

[0105] A 0.5-g portion of the polyorganosiloxane particles recovered from the latex by drying was immersed in 80 ml of folluene at 23 °C for 24 hours and, after 60 minutes of centrifugation at 12,000 rpm, the weight fraction (%) of the toluene-insoluble matter in the polyorganosiloxane particles was determined.

[Acetone-insoluble matter content]

[0106] One gram of the graft copolymer was immersed in 80 ml of acetone at 23 °C for 48 hours. Then, after 10

minutes of centrifugation at 18, 000 rpm, the sediment was weighed, and the acetone-insoluble matter content (%) was calculated

[Volume average particle diameter]

[0107] The volume average particle diameters of the seed polymer, polyorganositoxane particles and graft copolymer swere determined each in a latex form. Using LEED & NORTHRUP INSTRUMENTS MIGROTIAC UPA as the measuring apparatus, the volume average particle diameter (gurl) and the coefficient of variation in particle diameter distribution (standard deviation/volume average particle diameter (%)) were measured by the light scattering method.

[Impact resistance]

[0108] The evaluation was made by carrying out the Izod type test at -10 °C or 23 °C using notched 1/8 inch bars according to ASTM Date.

[Flame retardancy]

[0109] The evaluation was made by carrying out the test V according to UL 94.

20 [Hydrophilicity]

[0110] Latex of seed polymer was weighed out in a beaker in such an amount that solid content of the seed polymer was about 5 g. The latex was completely dried in a drier at 120 °C, and was weighed precisely. A 100 g portion of water was added to the dry seed polymer, stirred, with a stirrer, at 23 °C for 1 hour, and then filtered through filter paper. The filtrate was dried in a drier at 120 °C to recover water-soluble matters, and then the water-soluble matters were weighed precisely. The extraction rate of the water-soluble matters in dry seed polymer was calculated.

[Swelling capacity]

Journal of the seed polymer in a latex form was measured by MICROTRAC UPA. A emulsified liquid obtained by mixing organosiloxane (octamethylcycloterasiloxane), in an amount of 50 times (by weight) that of the seed polymer in a dry state, and 0.1% (by weight) equeous solution of Emal 2F (product of Kap) was incorporated into 5% (by weight) latex of the seed polymer and mixed. After one-hour stirring at 23 °C, particle diameters were measured by MICROTRAC UPA. The ratle of swelling by volume was calculated by the following equation;

(The rate of swelling by volume) =

{(Particle diameter after swelling measured in latex) /

(Particle diameter before swelling measured in latex))3 -1

First aspect of the invention

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45 (Reference Example 1) Production of polyorganosiloxane particles (S-1)

[0112] An emulsion was prepared by stirring an aqueous solution composed of the following components at 10,000 rpm for 5 minutes using a Homomico

	Amount (parts)
Component	251
Pure water	1.0
Sodium dodecylbenzenesulfonate (SDBS)	95
Octamethylcyclotetrasiloxane (D4) Mercaptopropyldimethoxymethylsilane (MPDS)	5
Mercaptopropyldimetric	

[0113] A 5-necked flask equipped with a stirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and thermometer was charged with the above errulsion all at once. With stirring the system, a 10% aqueous solution of do-

decy/benzenesulfonic acid (DSA) (1 part as solid content) was added, and the temperature was raised to 80 °C over about 40 minutes, and the reaction was allowed to proceed at 80 °C for 6 hours. Then, the mixture was cooled to 25 °C and, after 20 hours of standing, the plf of the system was returned to 6.5 with sodium hydroxide to terminate the °C and, after 20 hours of standing, one planting polygranositoxane particles (S-1) was thus obtained. The degree of conversion, and the volume average particle diameter and tolened-insoluble matter content of the polygranositoxane particle latex were determined. The results are shown in Table 1.

(Reference Example 2) Production of polyorganosiloxane particles (S-2)

[0114] A 5-necked flask equipped with a stirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and thermometer was charged with the following components.

Component	Amount (parts)
Pure water	189
SDBS	1.2

[0115] Then, while substituting the system atmosphere with nitrogen, the temperature was raised to 70°C and an aqueous solution composed of 1 part of pure water and 0.02 parts of potassium persulfate (KPS) was added. Then, a mixture composed of

Component	Amount (parts)
Styrene (St)	0.7
Butyl methacrylate (BMA)	1.3

was added all at once, and the polymerization was driven to completion by stirring for 1 hour to give an St-BMA copolymer latex. The degree of conversion in polymerization was 99%. The solid content of the latex obtained was 1, 0%, and the volume average particle diameter was 0.01 µm. This time, the coefficient of variation was 39%. The solventinsoluble matter content of the St-BMA copplymer was 0%.

[0116] Separately, a mixture composed of the following components was stirred with Homomixer at 10,000 rpm for 5 minutes to give an emulsion of the polyorganosiloxane-forming components.

Pure water	t (parts)
SDBC	0
SDBS	.5
	5
D4 γ-Methacryloyloxypropyldimethoxymethylsilane	

[0117] Then, the SI-BMA copolymer-containing latex was maintained at 80°C, a 10% aqueous solution of DBSA [2] parts as solid content) was added to the system, the emulsion of the polyorganosiloxane-forming components menioned above was then added all at once and, after 6 hours of continued stirring, the system was cooled to 25°C and illowed to stand for 20 hours. Thereafter, the pH was adjusted to 8.4 with sodium hydroxide to terminate the polymerization. A latex containing polyorganosiloxane particles (S-2) was obtained. The degree of conversion in polymerization, and the volume average particle diameter and toluene-insoluble matter content of the polyorganosiloxane particle latex were determined. The results are shown in Table 1. As is estimable from the charge amounts and degree of conversion, the polyorganosiloxane particle latex were determined. The results are shown in Table 1. As is estimable from the charge amounts and degree of conversion, the polyorganosiloxane particle and 25% of the St-BMA copolymer component.

Table 1

	Reference Example 1	Reference Example 2
P-l	S-1	S-2
Polyorganosiloxane particles Degree of conversion of polyorganosiloxane component (%)	87	87
Average particle diameter (µm)	0.14	0.04
Average particle diameter (µ11)		

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Table 1 (continued)

	Reference Example 1	Reference Example 2
Coefficient of variation (%)	35	35
Toluene insoluble matter content (%)	0	0

(Examples 1 to 5 and Comparative Examples 1 to 4)

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[0118] A 5-necked flask equipped with a sirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and thermometer was charged with 300 parts of pure water, 0.2 parts of sodium formaldehyde sufloxylate (SFS), 0.01 parts of ethylenediaminetetraacetic acid disodium salts (EGTA), 0.0025 parts of inclif) suffate, and a mount specified in Table 2 of the polyorganositysane particles (A1). While stirring the system, the temperature was raised to 60 °C in a mixture of the monomer (s) (B) and radical polymerization initiator each nitrogen atmosphere. After arrival at 60 °C, a mixture of the monomer (s) (B) and radical polymerization initiator each specified in Table 2 and a specified in Table 2 was added at a once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Table 2 was added dropwise over 3 hours and, after completion of °C for 1 hour. Then, the monomer (C) specified in Table 2 was added dropwise over 3 hours and, after completion of °C for 1 hour. Then, the monomer (C) specified in Table 2 was added dropwise over 3 hours and, after completion of °C for 1 hour. Then, the monomer (C) specified in Table 2 was added dropwise over 3 hours and, after completion of °C for 1 hour. Then, the monomer (C) specified in Table 2 was added at a once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Table 2 was added dropwise over 3 hours and, after completion of °C for 1 hour. Then, the monomer (C) specified in Table 2 was added at once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Table 2 was added at once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Table 2 was added at once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Table 2 was added at once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Table 2 was added at once, and the system was stirred at 60 °C for 1 hour. Then, the monomer (C) specified in Ta

[0119] The latex was then diluted with pure water to a solid concentration of 15%, a 10% acqueous solution to cause otherwise. The consultated sturry was otherwise. The consultated stu

[0120] In Table 2, AIMA stands for allyl methacrylate, BA for butyl acrylate, MMA for methyl methacrylate, AN for acrylonitrile (they are all monomers). CHP for cumene hydroperoxide (radical polymerization initiator), and Polymer SP for solubility parameter of polymer of vinyl monomer (s) (C) (as determined by the method described herein).

Table 2

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	Γ		"	Examples		ľ				
	_	-	<u>'</u> -	Salidhilas	1		ဝိ	Comparative Examples	Examples	Γ
Pohormani	T	1	1	,	•	20	-	2	8	4
particles (A1) (parts)	<u>.</u>	2	2	٤	92	i	P.	۶	۾	. ,
골	2-5	1	ı	1	1	8		ı	1	8
	AIMA	60	2	-	60	6	1	-	15	ı
Vinyi monomer (B)	Æ	1	ı	1	0.5	1	1	7	ı	
(parcs)	a d	0.01	0.01	10.0	10:0	10.0	-	10.0	0.03	-
	MMA	┸	28	29	27	1	30	27	5	1
•	ů		ı	ı	ı	27.75	ı	١	ı	8
Vinyl monomer (C)	; ;	1	ı	ŀ	١	9.25	ı	١	ı	우
(S) III (A)	§ §	٠	900	0.06	90.0	0.08	90.0	90'0	90'0	0.08
Polymer SP	9	1	9.25	9.25	9.25	9:92	9.25	9.25	9.25	9.95
(Can) Can)	1	1	8	8	đ	g	١	66	66	i
Degree of conversion	<u> </u>	6 G	6 6	66	66	66	66	. 99	66	8
Acetone-insoluble matter	matter	—	8	8	88	25	45	80	92	9
Graft nolymer No.	و	Sg-1	SG-2	SG-3	SG-4	SG-5	SG'-1	2G'-2	SG'-3	SG-4
Grait polymor										

(Examples 6 to 11 and Comparative Examples 5 to 11)

Rendering polycarbonate resins flame-retardant

[0121] A polycarbonate resin (PC-1: Toughlon FN 2200A, product of IDEMTSU PETROCHEMICAL; or PC-2: Toughlon FN 1900A, product of IDEMTSU PETROCHEMICAL) and the polyorganosiloxane-based graft copolymer obtained in any of Examples 1 to 5 (SG-1 to SG-5) or the polyorganosiloxane-based graft copolymer obtained in any

- of Comparative Examples 1 to 4 (SC-1 to SC-4) were blended together according to the formulation shown in Table 3. The abbreviation PEP36 stands for a phosphorus-containing antioxidant (ADEKA STAB PEP36, product of ASAHI DENKA), and PTFE for polyteraflucroethylene (Polyflon FA-500, product of OAIKIN Industries).
- [0122] Pellets were produced by melting and kneading each compound at 270°C on a twin-screw extruder (TEX 44 SS, product of Japan Steal Works). The pellets obtained were molded into 1/8 inch test specimens for impact resistance evaluation and 1/16 inch specimens for fame retardancy evaluation using FANUC's FAS 100 B injection molding evaluation and 1/16 inch specimens for fame retardancy evaluation using FANUC's FAS 100 B injection molting evaluation machine set at a cylinder temperature of 280 °C. The test specimens obtained were evaluated by the evaluation methods described hereinahous.
- [0123] The results are shown in Table 3.

Table 3

\perp	Ι.	-	Examples	ples					Compar	Comparative Examples	amples		
4	- 1	-	-		2	=	5	٥	-	8	6	10	Ξ
<u> </u>	_	ş	ŝ	5	5	ı	ë	5	5	9	-	100	1
1	·	1	-	1	1	ē	1	ı	ı	1	8	ı	5
65		1	1	١	1	6	-	ı	1	1	1	ı	ı
1			ı	ı	ı	1	ı	ı	ı	ı	1	ı	ī
<u> </u>		1	6	1	1	ı	ı	ı	1	ı	1	ı	ı
<u> </u>		1	ı	က	ı	ı	ı	1	ı	1	ı	1	1
1		1	ı	ł	6	ī	ı	1	1	1	1	I	ı
1			ı	1	1	ı	6	١	1	1	1	1	ı
 			ı	1	1	1	ı	က	ı	1	1	1	ı
1	<u> </u>	Τ.	1	1	1	ı	ı	ı	က	1	1	1	1_
1			1	١	ı	ı	ı	1	t	က	6	<u>'</u>	1
0.3 0.3	0	l	0.3	0.3	0.3	0.3	0.3	0.3	0.3	0.3	63	6.0	6.9
0.5 0.	o.	0.5	0.5	0.5	0.5	0.5	0.5	0.5	65	0.5	0.5	0.5	5.
20	L"	8	40	25	8	52	22	8		22	9	5	8
ums (sec.)	_	ę	S.	ટ	ž	ટ્ટ	ş	ટ્ટ	ž	윋	٤	ş	ξes
-10°C 55 (kJ/m²)		8	65	9	55	35	45	8	8	\$	8	8	5
	l	l											

[0124] From Table 3, it is seen that the graft copolymer according to the first aspect of the invention can highly improve the flame retardancy-impact resistance balance of the polycarbonate resins.

(Example 12 and Comparative Examples 12 and 13)

Rendering a polycarbonate/polyethylene terephthalate mixed resin flame-retardant

[0125] PC-1, a polyethylene terephthalate resin (PET: BELLPET EFG-70, product of Kanebo Gohsen) and the polyorganosiloxane-based graft copolymer (SG-1) obtained in Example 1 or the polyorganosiloxane-based graft copolymer (SG'-1) obtained in Comparative Example 1 were blended together according to the formulation shown in Table 4. [0126] Pellets were produced by melting and kneading each compound at 270 °C on a twin-screw extruder (TEX 44 SS, product of Japan Steel Works). The pellets obtained were molded into 1/8 inch test specimens for impact resistance evaluation and 1/12 inch specimens for flame retardancy evaluation using FANUC's FAS 100 B injection molding machine set at a cylinder temperature of 260°C. The test specimens obtained were evaluated by the evaluation methods described hereinabove.

[0127] The results are shown in Table 4.

15			Table 4		
			Example 12	Comparative Example 12	Comparative Example 13
	Thermoplastic resin	PC-1	90	90	90
20	Pidode (esil)	PET	10	10	10
	Graft copolymer	SG-1	4	4	-
		SG-1	1 - 1	4	-
25	Antioxidant	PEP36	0.3	0.3	0.3
_	Dripping-preventing agent	PTFE	0.5	0.5	0.5
	Flame retardancy	Total combustion time	50	65	190
30	j	(see) Dripping	No	No	Yes
- 1	Impact resistance	23°C (kJ/m²)	75	60	41

[0128] From Table 4, it is seen that the graft copolymer according to the invention can highly improve the flame retardancy-impact resistance balance of the polycarbonate/polyethylene terephthalate resin.

Second aspect of the invention

(Examples 13 to 18)

[0129] Water (400 parts) and an amount (as solid content) given in Table 5 of sodium dodecylbenzenesulfonate (SDBS) were mixed up in a 5-necked flask equipped with a stirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and thermometer, then the temperature was raised to 50 °C and, after arrival of the liquid temperature at 50 °C officers. °C, nitrogen substitution was effected. Then, a mixture of 10 parts of butyl acrylate and 3 parts of tert-dodecylmercaptan was added. After 30 minutes, 0.01 parts (as solid content) of paramenthane hydroperoxide was added, and the polymerization was allowed to proceed for 1 hour. Thereafter, a mixture of 90 parts of butyl acrylate and 27 parts of tertification was allowed to proceed for 1 hour. The existing representation of the proceeding the process of process of process of the process o dodecylmercaptan was added continuously over 3 hours. The subsequent 2 hours of post-polymerization gave seed latex (seed 1 to seed 4). The weight average particle diameter, hydrophilicity and degree of swelling after synthesis were determined. The results are shown in Table 5.

[0130] A 5-necked flask equipped with a stirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and thermometer was charged with a seed polymer specified in Table 6 in an amount (as solid content) specified in Table 6. Then, an emulsion of polyorganosiloxane-forming components as separately prepared by stirring a mixture composed of 300 parts of water, 0.5 parts (as solid content) of SDBS, 95 parts of octamethylcyclotetrasiloxane and 5 parts of dimethylimethylsilylpropyl methacrylate (DSMA) at 7,000 rpm for 5 minutes using a Homomixer was added all at once. [0131] Then, a 10% aqueous solution of dodecy/benzenesulfonic acid (1 part as solid content) was added and, with stirring the system, the temperature was raised to 80 °C in a nitrogen atmosphere. After arrival at 80 °C, stirring was continued at 80 °C for 6 hours and, then, the mixture was cooled to 25 °C and allowed to stand for 20 hours. Thereafter,

the pH was adjusted to 8.4 with sodium hydroxide to stop the polymerization. Latex containing polyorganosiloxane particles was thus obtained.

diameter is shown in Table 6.

[0133] The later was then dikted with pure water to a solid concentration of 15%, a 25% aqueous solution of calcium choice (a parts as solid content) was added. The thus-obtained coagulated sturry was heated to 55°C, then cooled choice (a parts as solid content) was added. The thus-obtained coagulated sturry was heated to 55°C, then cooled to 50°C, and dehydrated and dried to give a polyorganositoxane-based graft copolymer in a provider form.

[0134] Then, a polycarbonate resim (Toughlon FN 2200A, product of IDEMITSU PETROCHEMICAL) and the above polycoganositosiane-based graft copolymer in powder form were blended together according to the formulation shown in Table 6. The dripping-preventing agent used was polyterafluoroethylene (Polyflon FA-500, product of DAIKIN In-in Table 6. The dripping-preventing agent used was polyterafluoroethylene (Polyflon FA-500, product of LOIKIN Indicatives), and the stabilizer used was a mixture of a phosphorus-containing antioxidant (ADEKA STAB PEP36, product of LOI Japan).

Of ASAHI DENKA) and a phenoic antioxidant (Topanol CA, product of ICI Japan).

[0135] Pellets were produced by melting and kneading the thus-obtained compound at 270 °C on a twin-screw extruder (TEX.44.SS, product of Japan Steel Works). The pellets obtained were molided into 1/8 inch test specimens for impact restance evaluation and 1/16 inch specimens for flame retardancy evaluation using FANUCS FAS 100 B injection molding machine set at a cylinder temperature of 280 °C. The test specimens obtained were evaluated by the evaluation methods described hereinabove. The results of the impact resistance and flame retardancy evaluations of the moldings are shown in 2 to 16 6.

(Examples 19 and 20)

[0136] A 5-necked flask equipped with a stirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and thermometer was charged with a seed polymer specified in Table 6. In an amount (as solid content) specified in Table 6. Then, an emulsion of polyorganosiloxane-forming components as separately prepared by stirring a mixture composed of 300 parts of water, 0.5 parts (as solid content) of SDBS, 95 parts of octamethylcyclotetrasiloxane and 5 parts of 300 parts of water, 0.5 parts (as solid content) of SDBS, 95 parts of octamethylcyclotetrasiloxane and 5 parts of 100 parts of water, 0.5 parts (as solid content) and 100 parts of water was added all at once. In 10137] Then, a 10% aqueous solution of dode-cytheursensulfonic acid (1 part as solid content) was added and, with stirring the system, the temperature was raised to 80 °C in a nitrogen atmosphere. After arrival at 80 °C, stirring was routined at 80 °C for 8 hours and, then, the mixture was cooled to 25 °C and allowed to stand for 20 hours. Thereafter, the pH was adjusted to 6.4 with solitum hydroxide to terminate the polymerization. Latex containing polyorganositoxane

particles was thus obtained. The substance of the particle was thus obtained. (1918)

Then, a 5-necked flask equipped with a stirrer, reflux condenser, nitrogen inlet, inlet for adding monomer and (1918)

Then, a 5-necked flask equipped with a stirrer, reflux condenser, a lattice particle above polyorganositoxare thermoenter was charged with 240 parts of pure water and 70 parts (as solid content) of the above polyorganositoxare particles. With stirring the system, the temperature was raised to 40°C in a nitrogen amosphere. After arrival at 40°C, 21°C and 40°C oddim formatledly self-victory strength of the strength

[0139] The late was then diluted with pure water to a solid concentration of 15%, a 25% aqueous solution of calcium chloride (4 parts as solid content) was maded. The thus-obtained coagulated slury was heated to 85°C, then cooled to 50°C, and dehydrated and dried to give a polyorganosioxane-based graft copolymer in a powder form.

[0139] The late was then diluted with pure water to a solid concentration of 15%, a 25% aqueous solution of calcium chloride. The thus-obtained coagulated slury was heated to 85°C, then cooled to 50°C, and dehydrated and dried to give a polyorganosioxane-based graft copolymer in a powder form.

[0140] Then, a polycarbonate resin (Toughton FN 1900A, product of IDEMITSU PETROCHEMICAL) and the above polycrganosaloxane-based graft copolymer in Powder form were blended together according to the formulation shown in Table 6. The dripping-preventing agent used was polytetrafluoroethylene (Polyflon FA-500, product of DAIKIN Industries).

[0141] Pellets were produced by melting and kneading the thus-obtained compound at 270 °C on a twin-screw extractor (TEX 44 SS, product of Japan Steel Works). The pellets obtained were molded that 1/8 inch test specimens for impact resistance evaluation and 1/16 inch specimens for flame retardancy evaluation using FANUCS FAS 100 B injection molding machine set at a cylinder temperature of 280 °C. The test specimens obtained were evaluated by the evaluation methods desorbled hereinabove.

of the moldings are shown in Table 6.

(Comparative Example 14)

[0142] Formulation, molding and evaluations were carried out in the same manner as in Examples 13 to 18 except that the polyorganosiloxane-based graft copolymer was not added in formulating the polycarbonate resin composition. The results are shown in Table 6.

(Comparative Example 15)

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[0143] Synthesis, coagulation, heat treatment, dehydration/drying/powder formation, formulation, molding and evaluations were carried out in the same manner as in Examples 13 to 18 except that no seed polymer was added in polymerizing the latex containing polyorganosiloxane particles. The results are shown in Table 6.

5 (Comparative Example 16)

[0144] Formulation, moking and evaluations were carried out in the same manner as in Examples 19 and 20 except that the polycogeneous coare-based graft copolymer was not added in formulating the polycarbonate resin compositions. The results are shown in Table 6.

(Comparative Example 17)

[0145] Synthesis, coagulation, heat treatment, dehydration/drying/powder formation, formulation, molding and evaluations were carried out in the same manner as in Examples 19 to 20 except that no seed polymer was added in polymerizing the latex containing polyorganosiloxane particles. The results are shown in Table 6.

Table 5

	Seed 1	Seed 2	Seed 3	Seed 4
Sodium dodecylbenzenesulfonate (phr)	8	4	2	1
Butyl acrylate (phr)	100	100	100	100
tert- Dodecylmercaptan (phr)	30	30	30	30
Average particle diameter (µm)	0.04	0.06	0.08	0.09
Hydrophilicity (%)	80	80	80	80
Swelling capacity (times)	10	8	6	5

Table 6

	ib.	16	6	_		-		Т	7	छ।	4	-		Т	\neg	
	l		1	100	0.3	1	6	1	1	MPrDMS	ALMA	MMA	0.2	8	S.	7
	Comparative Examples		16	90	0.3	1	-	1	1	1	1	1	1	20	130	Not-V
	mparative	ľ	٩	8	0.5	9.0	69	-	-	DSMA	1	MMA	0.2	8	6	2
	õ		•	8	9.5	9.0	1	1	1	1	1	1	1	10	180	Not-V
-		8	1	8	0.3	1	3	Seed 1 Seed 2	1	DSMA MPrDMS MPrDMS	ALMA	MMA	0.15	20	45	٩
1		10	1	8	0.3	1	8	Seed 1	-	MPrDMS	ALMA	MMA	0.18	20	99	٩ >
1		18	1	3	0.5	9.0	3	Seed 4	2	DSMA	ı	MMA	0.22	22	5	오 >
		4	٤	3	0.5	9.0	3	Seed 3	7	DSMA	1	MMA	120	S	\$?
Syamples		9	5	1	0.5	9.0	က	Seed 2	2	DSMA	1	MMA	8	8		٩ >
	1	15	100	1	0.5	9.0	65	Seed 2 Seed 1 Seed 2 Seed 3 Seed 4	2	DSMA	1	MMA	0.2	20	各	٩ >
	:	•	5	t	0.5	9.0	67	Seed 2	-	DSMA		MMA	0.25	8	8	?
	5	2	8	t	0.5	9.0	65	Seed 1	-	DSMA	1	MMA	0.23	ន	6	7
	_		Perycarbonate (parts)	Oripping-preventing agent	(parts)	Stabilizer (parts)	Flame retardant addition level (parts)	Seed polymer	Seed amount (parts)	Conf. conselinking agent	The state of the s	Turnetters graff	Average particle diameter	(kJ/m²)	Total combustion	UL 94 V grading
			Leiycar	Oripping-pre	9	Stabili	Flame ret	Ses	Sond	4			Average	Impact	resistance	retardancy
		H	_		elur	.صد	1		1	nebr	ste1	ems[1	esi	tenst	Charac

INDUSTRIAL APPLICABILITY

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[0146] The invention can provide a flame retardant capable of giving thermoplastic resin compositions excellent in

flame retardancy-impact resistance balance when added to thermoplastic resins. Furthermore, thermoplastic resin compositions excellent in flame retardancy-impact resistance can be provided when the flame retardant is incorporated in thermoplastic resins.

Claims

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A polyorganosiloxane-containing graft copolymer

which is obtainable by polymerizing 0.5 to 10 parts by weight of a vinyl monomer (B) comprising 100 to 50% by weight of a polyfunctional monomer (b-1) containing two or more polymerizable unsaturated bonds and 0 to 50% by weight of another copolymerizable monomer (b-2), in the presence of 40 to 90 parts (as solid content) by weight of polyorganosiloxane particles (A1).

followed by further polymerization of 5 to 50 parts by weight of a vinyl monomer (C),

with the sum of (A1), (B) and (C) being 100 parts by weight.

- 2. The polyorganosiloxane-containing graft copolymer according to Claim 1, wherein the polyorganosiloxane particles (A1) have a volume average particle diameter of 0.008 to 0.6 μm.
- 3. The polyorganosiloxane-containing graft copolymer according to Claim 1 or 2, wherein the vinyl monomer (C) gives a polymer thereof having a solubility parameter of 9.15 to 10.15 (cal/
 - 4. The polyorganosiloxane-containing graft copolymer according to any of Claims 1 to 3, wherein the polyorganosiloxane particles (A1) are in a latex form.
- 5. The polyorganosiloxane-containing graft copolymer according to any of Claims 1 to 4,
 - wherein the vinyl monomer (C) comprises at least one monomer selected from the group consisting of aromatic vinyl monomers, vinyl cyanide monomers, (meth)acrylate ester monomers and carboxyl group-containing vinyl monomers.
- 6. A flame retardant

which comprises the polyorganosiloxane-containing graft copolymer according to Claim 1.

7. A flame retardant resin composition

which comprises 0.1 to 30 parts by weight, per 100 parts by weight of a thermoplastic resin, of the flame relardant according to Claim 6 as incorporated in the thermoplastic resin.

8. The flame retardant resin composition according to Claim 7, wherein the thermoplastic resin is a polycarbonate resin.

9. A polyorganosiloxane-containing graft copolymer

which is obtainable by polymerizing 0 to 10 parts by weight of a vinyl monomer (B) comprising 100 to 50% by weight of a polyfunctional monomer (b-1) containing two or more polymerizable unsaturated bonds and 0 to 50% by weight of another copolymerizable monomer (b-2), in the presence of 30 to 95 parts by weight (as solid content) of a polyorganosiloxane (A2) in a latex form obtainable by seed polymerization using, as a seed polymer,

a hydrophilic polymer capable of swelling in the corresponding organosiloxane, followed by further polymerizing 5 to 70 parts by weight of a vinyl monomer (C),

with the sum of (A2), (B) and (C) being 100 parts by weight.

10. The polyorganosiloxane-containing graft copolymer according to Claim 9,

wherein the seed polymer has such a degree of hydrophilicity that the extraction rate of water-soluble components in dry seed polymer is 10 to 100% by weight, as determined after adding water, in an amount of 20 weighttimes that of the seed polymer in a dry state, to the dry seed polymer, followed by stirring at 23 °C for 1 hour and wherein the seed polymer shows such a degree of swelling in the organosiloxane that the rate of swelling by volume as determined from the ratio between the latex particle diameter after stirring and the latex particle diameter before stirring is 3 to 50 times after adding the organosiloxane, in an amount 50 times by weight that of the dry seed polymer, to the seed polymer latex, followed by stirring at 23 °C for 1 hour.

11. The polyorganosiloxane-containing graft copolymer according to Claim 9,

wherein the seed polymer has such a degree of hydrophilicity that the extraction rate of water-soluble components in dry seed polymer is 50 to 100% by weight, as determined after adding water, in an amount of 20 weighttimes that of the seed polymer in a dry state, to the dry seed polymer, followed by stirring at 23 °C for 1 hour and wherein the seed polymer shows such a degree of swelling in the organosiloxane that the rate of swelling by volume as determined from the ratio between the latex particle diameter after stirring and the latex particle diameter before stirring is 3 to 15 times after adding the organosiloxane, in an amount 50 weight-times that of the dry seed polymer, to the seed polymer latex, followed by stirring at 23 °C for 1 hour.

12. The polyorganosiloxane-containing graft copolymer according to any of Claims 9 to 11,

Wherein the vinyl monomer (C) comprises at least one monomer selected from the group consisting of aromatic vinyl monomers, vinyl cyanide monomers, (meth)acrylate ester monomers and carboxyl group-containing vinyl monomers.

13. A flame retardant

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which comprises the polyorganosiloxane-containing graft copolymer according to Claim 9.

14. A resin composition excellent in impact resistance and flame retardancy

which comprises a thermoplastic resin and the flame retardant according to Claim 13 as incorporated therein in an amount of 0.1 to 30 parts by weight per 100 parts by weight of the thermoplastic resin.

15. The resin composition according to Claim 14,

wherein the thermoplastic resin is a polycarbonate resin.

	INTERNATIONAL SEARCH REPO	RT [pcT/JP03/01	622
A. CLA	ASSIFICATION OF SUBJECT MATTER L.Cl ⁷ CO8F283/12, CO8L51/08, 69	/00		
According	g to International Patent Classification (IPC) or to both n	national classification and	IPC .	
B. FIRI	DS SEARCHED			
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Document	ation searched other than minimum documentation to the	e extent that such docum	ents are included in the field	s searched
	data base consulted during the international search (nan	ne of data base and, when	s practicable, search lerms t	sed)
Category*	Citation of document, with indication, where a	romoriate, of the relevant	passages Relevan	to claim No.
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A	JP 7-33836 A (Mitsubishi Ray 03 February, 1995 (03.02.95) All references (Family: none	,	:	l-15
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Furthe	r documents are listed in the continuation of Box C.	See patent family	annex.	
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